APPENDIX B

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Introduction to Colloid and Surface Chemistry

Third Edition

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dan exchange

kinds of countarions are present, and can be represented by the equation Ion exchange involves an electric double layer situation in which two

RA + B = RB + A

concentration and charge number are of primary importance. R may be a cation exchanger (fixed negatively charged groups, such as -SO3 or position in the electric double layer around R, and, in this respect, where R is a charged porous solid. Counter-ions A and B compete for as -NHJ). A range of highly porous synthetic cation and anion exchange fairly rapid ion exchange. rasins are available commercially. The porcestry of the restn facilitates -COO) or an anion exchanger (fixed positively charged groups, such

of water and the 'defonisation' of water. The most important applications of ion exchange are the softening

resin, thus softening the water. Regeneration of the resin is effected by of a catlon exchange resin usually saturated with sodium counter-ions passing a strong-solution of sodium chloride through the column. ions in the water exchange with the singly charged sodium lons in the The doubly charged (and, therefore, more strongly adsorbed) calcium In the first of these processes, hard water is passed through a column

exchange with the cations and anions in the water sample and combine of a mixed ion exchange resin. These hydrogen and hydroxyl ions exchange resin saturated with hydroxyl ions are used, often in the form A cation exchange resh saturated with hydrogen lons and an anion The 'deionisation' of water involves both anion and cation exchange

the separation of the rare earths is usually achieved by ceition exchange followed by elution of their complexes with citric acid. Ion exchange has many preparative and analytical uses; for example,

ELECTROKINETIC PHENOMENA¹⁰⁷

which arise when attempts are made to shear off the mobile part of the Electrokinetic is the general description applied to four phenomena electric double layer from a charged surface.

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surface (plus attached material) tends to move in the appropriate direcrelative to each other. surface and the diffuse part of the double layer are made to move thus causing its flow. Conversely, an electric field is created if the charged migration in the opposite direction, carrying solvent along with them, tion, while the ions in the mobile part of the double layer show a not force is exerted on both parts of the electric double layer. The charged If an electric field is applied tangentially along a charged surface, a

The four electroidnetic phenomena are as follows:

- (1) Electrophoresis the movement of a charged surface plus alteshed material (i.e., dissolved or suspended material) relative to stationary liquid by an applied electric field.
- ন্ত Cleano-osmosis — the movement of liquid relative to a stallonary electric field (i.e. the complement of electrophoresis). The pressure necessary to counterbalance electro-asmotic flow is termed the charged suches (e.g. a capillary or porous plug) by an applied electro-osmotic pressure.
- (3) Streaming potential —the electric field which is created whon liquid is made to flow along a stationary charged surface (i.e. the opposite of electro-osmosis).
- \mathfrak{E} Sedimentation potential — the electric field which is created when of electrophoresis). charged particles move relative to stationary liquid (i.e. the opposite

owing to experimental difficulties. electrokinetic phenomena and has been studied extensively in its been studied to a moderate extent and sedimentation potential rurely various forms, whereas electro-osmosis and streeming potential have Electrophoresis has the greatest practical applicability of these

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tion of colloidal material in an electric field. A number of techniques have been developed for studying the inigra-

·Particle (microscopo) electrophoresis

suspension or emulsion containing microscopically visible particles or If the material under investigation is in the form of a reasonably stable

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